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(54) Title: IMMUNIZING AGAINST HIV INFECTION

(57) Abstract: A virus neutralizing level of antibodies to a primary HTV isolate is generated in a host by a prime-boost administration of antigents. The primary antigen is a DNA molecule encoding an envelop glycoprotein of a primary isolate of HTV-1 while the boosting antigen is either a non-infectious, non-replicating HTV-like particle having the envelope glycoprotein of a primary isolate of HTV-1 or an attenuated viral vector expressing an envelope glycoprotein of a primary isolate of HTV-1.

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TITLE OF INVENTION IMMUNIZING AGAINST HIV INFECTION

FIELD OF THE INVENTION

[0001] The present invention relates to the field of immunology and, in particular, to methods and compositions for immunizing a host against infection with HIV.

BACKGROUND OF THE INVENTION

[0002] Human immunodeficiency virus is a human retrovirus and is the etiological agent of acquired immunodeficiency syndrome (AIDS). It is estimated that more than 33 million people have been infected with HIV world-wide as of December 1999 (Ref 1- various references are referred to in parenthesis to more fully describe the state of the art to which this invention pertains. Full bibliographic information for each citation is found at the end of the specification, immediately preceding the claims. The disclosure of these references are hereby incorporated by reference into the present disclosure).

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[0003] As the HIV epidemic continues to spread world wide, the need for an effective vaccine remains urgent. Efforts to develop such a vaccine have been hampered by several factors three of which are: (a) the extraordinary ability of the virus to mutate; (b) inability of most known specificities of anti-HIV antibodies to neutralise HIV primary isolates consistently; and (c) lack of understanding of the correlates of protective immunity to HIV infection. Over the last 10 years, several candidate HIV vaccines have been tested in primates for their immunoprotective abilities (Ref. 2). These studies suggest that both neutralising antibodies and cell-mediated immunity play a role in conferring sterilizing immunity and preventing progression towards disease (Ref 3, 4). While the correlates for immune protection against HIV-1 infection are currently unknown, an effective HIV vaccine should elicit both strong neutralising antibody and cytotoxic T lymphocyte (CTL) responses.

[0004] Envelope subunit vaccines have been shown to induce high titred humoral responses, but were inefficient in eliciting CTL responses (Ref 5). Live recombinant pox vectors have been shown to elicit very potent CTL responses, however these vectors were ineffective for generating a significant antibody

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response (Ref 6). In attempts to combine the two immunization types, several clinical trials involved a prime-boost strategy, consisting of initial viral vector immunization followed by boosts with recombinant HIV-1 envelope subunits (Ref 7, 8), have led to limited success with respect to CTL responses. Other vaccine approaches have used non-infectious, non-replicating, immunogenic virus-like particles (VLP) for immunising against HIV infection (Ref 9, 10). This type of immunogen has lead to the generation of neutralizing antibodies to a laboratory HIV-1 strain (Ref 10).

[0005] A prime-boost approach has been investigated using non-infectious VLPs to enhance HIV-specific CTL responses in mice primed with recombinant canarypox vector vCP205 encoding HIV-1gp 120 (MN strain) (Ref 11). This study showed that VLPs could boost the CTL response to the canarypox vector.

[0006] Recently, a study showing the induction of neutralizing antibodies to a HIV-1 primary isolate in chimpanzees has been reported (Ref 12). In this study, recombinant adenovirus expressing gp160 was used as the priming agent and recombinant gp120 protein was used to boost the monkeys.

[0007] There is still a need for vaccines and immunization regimes to induce both a strong CTL response as well as neutralizing antibodies to HIV primary isolates.

SUMMARY OF THE INVENTION

[0008] In accordance with one aspect of the present invention, there is provided a method for generating, in a host, particularly a human host, a virus neutralizing level of antibodies to a primary HIV isolate, comprising at least one administration of a priming antigen to the host, wherein the priming antigen comprises a DNA molecule encoding an envelope glycoprotein of a primary isolate of HIV, resting the host for at least one specific resting period to provide for clonal expansion of an HIV antigen specific population of precursor B-cells therein to provide a primed host, and at least one administration of a boosting antigen to the primed host to provide said neutralizing levels of antibodies, wherein the boosting antigen is selected from the group consisting of a non-infectious, non-replicating, immunogenic HIV-like particle having at least part of the envelope glycoprotein of a primary isolate of HIV and an attenuated viral

vector expressing at least part of an envelope glycoprotein of a primary isolate of HIV.

[0009] The primary HIV isolate may be an HIV-1 isolate including from the clade B HIV-1 clinical isolate HIV- $1_{\rm Bx08}$, although any other primary HIV-1 isolate may be employed in the immunization procedures of the invention.

[0010] The DNA molecule encoding the envelope glycoprotein of a primary isolate of HIV may be contained in a plasmid vector under the control of a heterologous promoter, preferably a cytomegalovirus promoter, for expression of the envelope glycoprotein in the host, which may be a human host.

10 [0011] The vector utilized for DNA molecule immunization is novel and constitutes a further aspect of the present invention. Preferably, the vector has the identifying characteristics of pCMV3Bx08 shown in Figure 2, such identifying characteristics being the nucleic acid segments and restriction sites identified in Figure 2.

15 [0012] A priming administration of antigen may be effected in a single or in multiple administrations of the priming antigen. In the latter case, the at least one specific resting period to permit clonal expression of HIV antigen-specific population precursor B-cells may be effected after each priming administration. The at least one specific resting period may be between about 2 and 12 about 20 months.

[0013] In the embodiment where the boosting antigen is a non-infectious, non-replicating, immunogenic HIV-like particle, such particle may comprise an assembly of:

(i) an env gene product,

a gag gene product

25 (ii) a pol gene product, and

(iii)

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with the particle being encoded by a modified HIV genome deficient in long terminal repeats (LTRs) and containing gag, pol and env in their natural genomic arrangement. Such particles and the manufacture thereof are described in US Patent No. 5,439,809, assigned to the assignee hereof and the disclosure of which is incorporated herein by reference. Such particles can include mutations in gag

and pol to further reduce potential infectivity, as more fully described in United

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States Patent No. 6,080,408, assigned to the assignee hereof and the disclosure of which is incorporated herein by reference (WO 96/06177). In a preferred embodiment, the *env* gene is that from primary isolate BX08. The *gag* gene and *pol* gene may be those from the same primary isolate or may be chosen from those of other HTV-1 isolates, which may be primary isolates.

[0014] The non-infectious, non-replicating, immunogenic HIV-like particle may be administered in conjunction with an adjuvant. Any suitable adjuvant may be used, such as QS21, DC-chol, RIBI or Alum.

[0015] Such non-infectious, non-replicating, immunogenic HIV particle may be formed by expression from a suitable vector in mammalian cells. In accordance with an additional aspect of this invention, there is provided a vector comprising a modified HIV-genome deficient in long terminal repeats and a heterologous promoter operatively connected to said genome for expression of said genome in mammalian cells to produce the non-infectious, non-replicating and immunogenic particle, wherein at least the *env* gene of the modified HIV-genome is that from a primary isolate of HIV. The *gag* and *pol* genes of the modified HIV genome may be those from the same primary isolate or those from another isolate, which may be a primary isolate.

[0016] The vector preferably is a plasmid vector while the primary isolate preferably is BX08. The promoter may be the metallothionein promoter. The vector preferably has the identifying characteristics of plasmid p133B1 shown in Figure 3, such identifying characteristics being the nucleotide segments and restriction sites identified in Figure 3.

viral vector, the attenuated viral vector may be an attenuated avipox virus vector, particularly the attenuated canary poxvirus ALVAC. The attenuated viral vectors used herein form another aspect of the invention. The attenuated viral vector may contain a modified HIV genome deficient in long terminal repeats (LTRs), wherein at least the *env* gene is that from primary isolate BX08. The *gag* and *pol* genes of the modified genome may be those from the same primary isolate or may be chosen from other HIV isolate.

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[0018] The attenuated canarypox virus-based vector ALVAC is a plaquecloned derivative of the licensed canarypox vaccine, Kanapox, and is described in reference 19. The attenuated canary pox vector preferably has the identifying characteristics of vCP1579 shown in Figure 4, such identifying characteristics being the nucleic acid segments and restriction sites identified in Figure 4.

[0019] The at least one administration of a boosting antigen may be effected in a single administration or at least two administration of the boosting antigen.

[0020] The invention further includes compositions comprising the immunogens as provided herein and their use in the manufacture and formulation of immunogenic compositions including vaccines.

BRIEF DESCRIPTION OF THE DRAWINGS

[0021] The present invention will be further understood from the following description with reference to the drawings, in which:

Figure 1 shows the details of the elements of plasmid pCMVgDtat vprBx08.

Figure 2 shows the details of the elements of plasmid pCMV3Bx08.

Figure 3 shows the details of the elements of plasmid p133B1.

Figure 4 shows the details of the insertions into ALVAC (2) to provide vector vCP1579.

Figures 5A and 5B contain a representation in time-line form of the immunization regime used wherein the study groups are described in Table 1. The numbers below the lines refer to weeks.

Figure 6 shows the immunoreactivity to HIV-1 antigens of the serum diluted 1:100 from the macaques immunized with the various preparations as described in Table 1.

Figure 7 shows the immunoreactivity to HIV-1 antigens of the serum diluted 1:1000 from the macaques immunized with the various preparations as described in Table 1.

Figure 8 shows the details of the elements of pMPC6H6K3E3. Figure 9 shows the details of the elements of pMPC5H6PN.

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Figure 10 shows the details of the elements of pHIV76.

Figure 11 shows the nucleotide sequence (SEQ ID NO: 1) for the H6/HIV Pol/Nef epitope cassette in the ALVAC C5 site of vCP1579.

Figure 12 contains the nucleotide sequence of C6 region (coding strand SEQ ID NO: 16, complementary strand SEQ ID NO: 17, K3L amino acid sequence SEQ ID NO: 18, E3L amino acid sequence SEQ ID NO: 19).

GENERAL DESCRIPTION OF INVENTION

[0022] As noted earlier, the present invention involves administration of HIV antigens to elicit virus-neutralizing levels of antibodies against a primary HIV isolate.

[0023] A DNA construct was prepared incorporating the *env* gene from the primary isolate Bx08 under the control of the cytomegalovirus promoter and the construct, pCMV3Bx08, is shown in Figure 2. The construct pCMV3Bx08 is derived from plasmid pCMVgDtat vprBx08 seen in Figure 1. The DNA construct pCMV3Bx08 was used in a priming immunization step to a host, macaque monkeys being the animal model chosen.

[0024] Following the priming immunization step, which may be effected in one or more administrations of the DNA construct, the host is allowed to rest to provide for clonal expression of an HIV antigen specific population of precursor B-cells therein to provide a primed host.

[0025] The boosting administration is effected either with a non-infectious, non-replicating, immunogenic HIV-like particle (VLP) or an attenuated viral vector.

[0026] For this purpose, a VLP expression plasmid was constructed containing a modified HIV genome lacking long terminal repeats in which the *env* gene is derived from primary isolate BX08, wherein the modified HIV genome is under the control of a metallothionein promoter. The construct, p133B1, shown in Figure 3, was used to effect expression in mammalian cells of the non-infectious, non-replicating, immunogenic HIV-like particules, in which the *env* gene product is that from the primary isolate BX08.

[0027] In the case of the attenuated virus vector, a recombinant attenuated canarypox virus vector was constructed to contain the env gene from primary

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isolate BX08. The viral vector vCP1579 (Figure 4) was prepared by a variety of manipulatious from plasmid pHIV76 (Figure 10), as shown described in detail below.

[0028] These products were utilized in a boosting administration to the primed macaques. The boosting administration may be effected in one or more immunizations. In a preferred aspect of the invention, the non-infectious, non-replicating immunogenic HIV-like particles may co-administered with the DNA construct in the priming administration and the DNA construct may be coadministered with the HIV-like particles in the boosting administration.

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10 [0029] Immunizations were effected in accordance with the procedure of the invention and the results obtained were compared with those obtained using other protocols according to the protocols set forth in Table 1. The immunization regimes used are shown as time lines in Figures 5A and 5B.

[0030] The results obtained following the various protocols showed that, in particular, a primary DNA vaccination in combination with a boost from either the VLP or the attenuated canarypox virus enhanced the levels of neutralizing antibodies, as indicated by the reduction of detectable p24 levels in cells infected with primary HIV isolates.

Biological Deposits

Certain vectors that are described and referred to herein have been [0031] 20 deposited with the American Type Culture Collection (ATCC) located at 10801 University Boulevard Manassas, Virginia 20110-2209, USA, pursuant the Budapest Treaty and prior to the filing of this application. Samples of the deposited vectors will become available to the public and all restrictions imposed or access to the deposits will be received upon grant of a patent based on this 25 United States patent application or the United States patent application in which they are described. In addition, the deposit will be replaced if viable samples cannot be dispensed by the Depository. The invention described and claimed herein is not limited in scope by the biological materials deposited, since the deposited embodiment is intended only as an illustration of the invention. Any 30 equivalent of similar vectors that contain nucleic acids which encode equivalent or

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similar antigens as described in this application are within the scope of the invention.

Deposit Summary

	Plasmid	ATCC	Deposit Date
5	pMT-HIV	40912	October 12, 1990
	pCMVgDtat vpr	209446	November 11, 1997
		EXAMPLES	

more complete understanding can be obtained by reference to the following specific Examples. These Examples are described solely for purposes of illustration and are not intended to limit the scope of the invention. Changes in form and substitution of equivalents are contemplated as circumstances may suggest or render expedient. Although specific terms have been employed herein, such terms are intended in a descriptive sense and not for purposes of limitation.

15 [0033] Example 1

[0034] This Example describes the construction of plasmid pCMV3BX08.

[0035] The plasmid, pCMV3BX08, contains sequence segments from various sources and the elements of construction are depicted in Figure 2.

[0036] The prokaryotic vector pBluescript SK (Stratagene) is the backbone of the plasmid pCMV3.BX08 and was modified by the replacement of the Amp^R with Kan^R gene and the deletion of the fl and the LacZ region. To achieve the desired modifications, the sequence between Ahdl (nucleotide 2,041) and Sacl (nucleotide 759) of pBluescript SK, which contains the Amp^R, fl origin and the LacZ, was deleted. A 1.2 kb Pstl fragment from the plasmid pUC-4K (Pharmacia) containing the Kan^R gene, was blunt end ligated to the Ahdl site of pBluescript SK in a counter-clockwise orientation relative to it's transcription. A 1.6 kb Sspl/Pstl DNA fragment containing the human cytomegalovirus immediate-early gene promotor, enhancer and intron A sequences (CMV) was ligated to the other end of the Kan^R gene so that the transcription from the CMV promoter proceeds in the clockwise orientation. A synthetic oligonucleotide segment containing translation initiation sequence and sequences encoding the human tissue plasminogen activator signal peptide (TPA) was used to link the

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CMV promotor and the sequences encoding the envelope gene of the primary isolate HIV-l_{BX08}.

The envelope gene from the HIV-1 primary isolate BX08 was [0037]isolated from the plasmid pCMVgDtat vprBx08 illustrated in Figure 1. The plasmid pCMVgDtat vpr Bx08 was derived from the deposited plasmid pCMVgDtat vpr, the construction of which is described in copending United States Patent Application No. 08/991,773 filed December 16, 1997, assigned to the assignee hereof and the disclosure of which is incorporated herein by reference, (WO 99/31250). The plasmid pCMVgDtat vpr Bx08 was derived by substituting the BX08 envelope sequence from clade B HIV-1 clinical isolate HIV-1_{BX08} for the modified HIV genome sequence present in pCMVgDtat vpr. Plasmid pCMVgDtat vpr Bx08 was restricted with the restriction enzyme Xho I and made blunt ended with Klenow treatment. A Not I partial digestion was then performed and the resulting 6.3 kb fragment containing the env gene was isolated. Plasmid pCMV3 (Invitrogen) was restricted with Bam HI and made blunt ended with Klenow treatment. The plasmid pCMV3 was then restricted with Not I and the resulting 4.4 kb fragment was isolated. The 6.3 and 4.4 kb fragments were

[0038] The pCMV3BX08 construct was introduced into HB101 competent cells according to manufacturer's recommendations (GibcoBRL). Correct molecular clones were identified by restriction and sequencing analysis and their expression of envelope glycoprotein was examined in transient transfections followed by Western blot analysis.

ligated together to produce plasmid pCMV3BX08 (Figure 2).

[0039] All DNAs used for immunizations were prepared using EndoFree Plasmid Kit (Qiagen). For intramuscular immunizations either 3 mg or 600 µg of pCMVBX08, in 100 µl PBS was injected.

[0040] Proviral DNA for clade B HIV-1 clinical isolate HIV-1_{BX08} originated at Transgene (Strasbourg, France) and was isolated from genomic DNA of cells infected with the virus.

30 [0041] <u>Example 2</u>

[0042] This Example describes the construction of plasmid p133B1.

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A Bx08 plasmid expression vector (p133B1, Figure 3) used to [0043] transfect the mammalian cells was engineered in several stages using pUC18 as the initial host plasmid. First, an 8.3-kbp fragment of HIV-1_{LAI} provirus encoding the gag, pol and env proteins was isolated. This fragment lacked the transcription regulatory elements and long terminal repeat elements from each end of the proviral genome to ensure the virus-like particles would be replication-This fragment was linked to an inducible human type IIA incompetent. metallothionein (MTIIA) promoter (Ref 13) and also to a Simian Virus 40 polyadenylation (polyA) addition/transcription termination sequence from plasmid pSV2dhfr (Ref 14). The modified fragment was then inserted into the pUC18 host vector. The resulting deposites expression construct, named pMT-HIV, was used to transfect into African green monkey kidney (Vero) and COS monkey kidney The procedure for obtaining pMT-HIV is further described in the cells. aforementioned US Patent No. 5,439,809. Both transfected cell lines produced non-replicating virus-like particles when induced with metal ions (Ref 15).

[0044] Two further modifications were made to the provinal DNA in pMT-HIV to provide additional safety features to protect human cells against recombination events with reverse-transcribed DNA:

- 1) inactivation of the RNA packaging sequences; and
- 20 2) deletion of a large section of the *pol* gene encoding reverse transcriptase and integrase.

[0045] To delete the first RNA packaging signal, part of the DNA corresponding to the untranslated leader sequence of the mRNA was replaced with synthetic DNA lacking a 25-bp motif corresponding to nucleotides 753-777 (the *psi* sequence). To inactivate the second RNA packaging signal, two adenosine residues within a *gag* gene zinc finger sequence were changed to thymidine residues. Each of these residue changes had the effect of replacing cysteine residues in a Cys-His array with a serine in the gene product.

[0046] The pol gene deletion was effected by replacing a 1.9-kbp fragment with synthetic DNA containing stop codons in all three reading frames. This prevented read-through translation of the residual integrase coding sequence on the 3' side of the deletion. The 1.9-kbp deletion in pol also eliminated the

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expression of reverse transcriptase and integrase enzymes. However, the deletion left intact the gene encoding the viral protease, which is both an immunogenic component of HIV-1 virus particles and allows the expression of particles with processed gag antigens closely resembling native virions (Ref 16). The protease also contains epitopes that are conserved across HIV-1 clades. The modifications described with respect to gag and pol genes are more fully described in the aforementioned United States Patent No. 6,080,408 (WO 96/06177).

[0047] Finally, the HIV-1_{LAI} env gene within pMT-HIV was replaced with that of HIV-1_{Bx08}. To effect this replacement, a 2440-bp fragment containing the env gene of Bx08 was amplified by polymerase chain reaction (PCR) from cells infected with this isolate. The PCR product was then used to replace the corresponding region present in pMT-HIV. However, the incoming fragment from HIV-1_{Bx08} was 125-bp shorter than the original HIV-1_{LAI} region owing to a deletion in the untranslated region between the env gene stop codon and the termination/polyA addition sequence. The resulting construct replaced all but eleven amino acid residues of the LAI envelope proteins gp120 and gp41. Of these eleven, only the first three differ between the LAI and Bx08 isolates, and these are all charge-conservative changes meaning the final expression vector (p133B1) encoded a nearly authentic HIV-1_{Bx08} env protein.

20 [0048] <u>Example 3</u>

[0049] This Example describes the production of HIV-like particles.

[0050] African green monkey kidney (Vero) cells were recovered and cultivated in Dulbecco's modified Eagle medium (DMEM) containing 10% v/v fetal bovine serum (FBS), referred to below as Complete Medium. At passage 141, the cells were transfected with p133B1 using the calcium phosphate method when at approximately 30% confluence. The cells were shocked with glycerol 8 hours after transfection. For this step, six 10-cm dishes containing approximately 3.0 x 10⁶ cells each in 10.0 mL of Complete Medium were prepared. Each dish received 25.0 µg of expression vector and 2.0 µg of plasmid pSV2neo (Ref 17). The pSV2neo contains a selectable marker gene conferring resistance to the antibiotic geneticin (G418). Two days after transfection, the cells from each dish

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were recovered by trypsinization and replated into twenty-five fresh dishes in Complete Medium supplemented with 0.5 mg/mL of G418.

[0051] In total, 394 colonies were isolated from the dishes using cloning cylinders. Each colony was recovered by trypsinization and divided into two cluster dish wells, one of the wells per clone was induced after reaching 50% to 90% confluence. Prior to induction, the wells were treated by replacing all the medium with fresh Complete Medium containing 10.0 μM 5-azacytidine. After incubating for between 18 hours and 22 hours, the medium was removed and replaced with fresh DMEM containing 0.2% v/v FBS, 2.0 μM CdCl₂ and 200.0 μM ZnCl₂. The wells were incubated for a further 20 hours to 24 hours at which time samples of the medium were removed and tested by p24 ELISA.

[0052] The twenty highest-producing clones, based on the p24 titre, were chosen and cells from the corresponding uninduced wells were sub-cultured into one T-25 and one T-150 flask per clone. Both flasks were grown to confluence.

The cells from the T-150 were recovered by trypsinization and cryopreserved at passage number 145. The cells from the T-25 were recovered by trypsinization every 3 days to 4 days and maintained up to passage 153. The cells were induced as above and samples retested by p24 ELISA at two different passages prior to passage 153.

20 [0053] The two highest p24 producers were chosen and were recovered by trypsinization every 3 days to 4 days up to passage 163. Samples from the clones were tested by p24 and gp120 ELISA from passage 158 and by p24 ELISA at passage 163, to assess clonal stability. The most suitable of these two cell lines, named 148 to 391, was chosen for further sub-cloning. The clone nomenclature defines the experiment number for this procedure, which was 148, and the number of the clone, which was number 391 of the original 394 isolated.

[0054] The vero cells were grown for approximately 100 h to 103 h and the medium was then replaced with growth medium containing 5-azacytidine. The bottles were then incubated for a further 20 h to 22 h, at which time the medium was replaced with serum-free medium containing CdCl₂ and ZnCl₂. The bottles were then incubated for 29 h to 31 h, at which time the medium was harvested, pooled and stored at 2°C to 8°C prior to purification.

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[0055] The next day after harvesting, the solution was clarified, concentrated and diafiltered against phosphate buffer. The concentrate was passed through a ceramic hydroxyapatite (type I) column and the run-through was collected. The run-through from two successive sublots was pooled together and pumped onto a sucrose density gradient in a continuous zonal ultracentrifuge rotor. Pseudovirion-containing fractions were collected and pooled. The pooled pseudovirion fractions were diafiltered against PBS containing 2.5% sucrose to reduce the sucrose content, concentrated and diafiltered again. The material was sterile filtered using a 0.2 μm filter. At this stage the materials was designated as a purified sub-lot and were stored at 2 to 8°C.

[0056] The adjuvants were prepared separately and filter sterilized before filling in single dose vials. QS21 was stored at -20°C.

[0057] Example 4

[0058] This Example describes the production of recombinant pox virus vCP1579.

[0059] Recombinant pox virus vCP1579 (Figure 4) contains the HIV-1 gag and protease genes derived from the HIV-1 IIIB isolate, the gp120 envelope sequences derived from the HIV-1 Bx08 isolate, and sequences encoding a polypeptide encompassing the known human CTL epitopes from HIV-1 Nef and Pol.

[0060] Recombinant vCP1579 (Figure 4) was generated by insertion of the vector modifying sequences from pMPC6H6K3E3 (Figure 8) encoding E3L and K3L into the C6 site of recombinant vCP1566 (Figure 4). Recombinant vCP1566 was generated by insertion of an expression cassette encoding a synthetic polypeptide containing Pol CTL epitopes and Nef CTL epitopes (Figure 11) and plasmid pMPC5H6PN (Figure 9) into vCP1453 at the insertion site known as C5. Recombinant vCP1453 was generated by co-insertion of genes encoding HIV-1 env and gag/protease gene products, plasmid pHIV76 (Figure 10), into the ALVAC genome at the insertion site known as C3.

The construction of recombinant pox vectors containing the E3L and K3L genes has been described in United States patent 6,004,777 issued Dec 21, 1999 to Tartaglia et al. and the recombinant pox vectors describing the

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insertion of HIV genes has been described in United States patent 5,766,598 issued June 16 1998 to Paoletti et al.

[0062] The locus designated C3 was used for the insertion of the HIV-1 env and gag gene sequences into the ALVAC(2) vector, and the locus designated as C5 was the insertion site for the sequences encoding the HIV-1 Nef and Pol CTL epitopes. By virtue of the C3 and C5 loci existing within the extensive inverted terminal repetitions (ITRs) of the virus genome (approximately 41 kbp), insertion into these loci results in the occurrence of two copies of the inserted HIV-1 sequences.

Briefly, expression cassette pHIV76 (Figure 10) was engineered in [0063] 10. the following manner. Plasmid p133B1 (Figure 3) containing the HIV-1Bx08 gp 160 gene was used as the starting plasmid. The 3'-end of the H6 promoter was cloned upstream of the gp160 gene and three poxvirus early transcription termination signal sequences (T₅NT) were modified. This was accomplished by cloning a 2,600 bp BamHI-digested PCR fragment, containing the 3'-end of the H6 promoter and the T₅NT-modified HIV-1 (BX08) gp160 gene, into the BamHI site of pBS-SK. This PCR fragment was generated from four overlapping PCR fragments (a 570 bp fragment, a 140 bp fragment, a 500 bp fragment and a 1,450 bp fragment) and the oligonucleotides, RW835 (5'-ATCATCATCGGATCC CGGGGTCGCGATATCCGTTAAGTTTGTATCGTAATGAAAGTGAAGGAC C-3' - SEQ ID NO: 2) and RW836 (5'-ATCATCATCGGATCCCGGGGTT ATAGCAAAGCCCTTTC-3' - SEQ ID NO: 3). The 570 bp PCR fragment, containing the 3'-end of the H6 promoter and the 5'-end of the gp160 gene, was generated from the plasmid, p133B1, with the oligonucleotides, RW835 (5'-ATC ATCATCGGATCCCGGGGTCGCGATATCCGTTAAGTTTGTATCGTAATG AAAGTGAAGGAGACC-3') and RW868 (5'-ATCAAGACTATAGAAGA GTGCATATTCTCTCTCATC-3'). The 140 bp PCR fragment, containing an interior portion of the gp160 gene, was generated from plasmid p133-B1 with the oligonucleotides, RW864 (5'-GCACTCTTCTATAGTCTTGATATAGTAC-3' -NO: 4) and RW865 (5'-AGCCGGGGCGCAGAAATGTATG SEQ GGAATTGGCAC-3' - SEQ ID NO: 5). The 500 bp PCR fragment, containing an interior portion of the gp160 gene, was generated from 133-3 with the

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oligonucleotides, RW866 (5'-ATACATTTCTGCGCCCCGGCTGGT TTTGCGATTC-3' - SEQ ID NO: 6) and RW867 (5'-GAAGAATTC CCCTCCACAATTAAAAC-3' - SEQ ID NO: 7). The 1,450 bp PCR fragment, containing the 3'-end of the gp160 gene, was generated from p133-B1 with the oligonucleotides, RW869 (5'-TGTGGAGGGGAATTCTTCTACTGTAATAC AACACAAC-3' - SEQ ID NO: 8) and RW836 (5'-ATCATCATCGGAT CCCGGGGTTATAGCAAAGCCCTTTC-3' - SEQ ID NO: 9). The 3'-end of the 570 bp PCR fragment overlaps the 5'-end of the 140 bp PCR fragment. The 3'-end of the 140 bp PCR fragment overlaps the 5'-end of the 500 bp PCR fragment.

fragment. The plasmid generated by this manipulation is called pRW997.

[0064] The sequence encoding gp41 was then replaced with the sequence encoding the gp160 transmembrane (TM) region. This modification was accomplished by cloning a 200 bp MfeI-HindIII-digested PCR fragment, containing the 3'-end of the gp120 gene and the TM sequence, into the 4,400 bp MfeI-HindIII fragment of pRW997. This PCR fragment was generated from two overlapping PCR fragments (a 170 bp fragment and a 125 bp fragment) with the

oligonucleotides, HIVP97 (5'-TAGTGGGAAAGAGATCTTCAGACC-3' - SEQ

ID NO: 10) and HIVP101 (5'-TTTTAAGCTTTTATCCCTGCCTAACT

20 CTATTCAC TAT-3' - SEQ ID NO: 11). The 170 bp PCR fragment was generated from pRW997 with the oligonucleotides, HIVP97 (5'-TAGTGGGAAAGAGATCTTCAGACC-3' - SEQ ID NO: 12) and HIVP100 (5'-CCTCCTACTATCATTATGAATATTCTTTTTTCTCTCTGCACCACTCT-3' - SEQ ID NO: 13). The 125 bp PCR fragment was generated from pRW997 with

the oligonucleotides, HIVP99 (5'-AGAGTGGTGCAGAGAGAAAAA AGAATATTCATAATGATAGTAGGAGGC-3' - SEQ ID NO: 14) and HIVP101 (5'-TTTTAAGCTTTTA TCCCTGCCTAACTCTATTCACTAT-3' - SEQ ID NO: 15). The plasmid generated by this manipulation is called pHIV71.

[0065] The H6-promoted gp120+TM gene was then cloned between C3 flanking arms, into a plasmid containing the I3L-promoted HIV1 gag/(pro) gene. This modification was accomplished by cloning the 1,600 bp Nrul-XhoI fragment of pHIV71, containing the H6-promoted gp120+TM gene, into the 8,200 bp Nrul-

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XhoI fragment of pHIV63. The plasmid generated by this manipulation is called pHIV76 (Figure 10). Plasmid pHIV76 was used in *in vivo* recombination experiments with ALVAC (CPpp) as rescue virus to yield vCP1453.

[0066] The sequence of the nef/pol regions is shown in Figure 12 and the E3L and K3L sequences are shown in Figure 13. To generate ALVAC(2)120(BX08)GNP (vCP1579), expression cassettes consisting of the promoter/HIV-1 gene combinations were subcloned into an ALVAC donor plasmid, which were then used to insert the expression cassettes into defined sites in the ALVAC genome by *in vitro* recombination as previously described (Ref 20).

[0067] <u>Example 5</u>

[0068] This Example describes the results of immunization regimes.

[0069] Groups of four animals (macaques) each were randomly assigned to seven vaccine groups as illustrated in Table 1. In this Table, "BX08 DNA" refers to pCMV3BX08, prepared as described in Example 1, "BX08 VLP" refers to the pseudovirions produced by expression vector p133B1 in Vero cells, as described in Example 3, and "ALVAC(2) BX08" refers vCP1579, prepared as described in Example 4. Reference (pre-bleed) sera were sampled at -6 and -2 weeks pre-vaccination. Primary immunizations with the various vaccines were given on weeks 0 and 4 with boosts on weeks 24 and 44 (Figures 5A, 5B). The vaccines were immunized intramuscularly into one quadricep of each macaque monkey.

[0070] Sera were prepared from whole-blood using SST collection tubes and analyzed using commercially available HIV-1 western blots. Groups 1, 2 and 7 showed low levels of anti-Env antibodies after the first boost (Figures 6 and 7). Based on ELISA values, the anti-env antibody levels were below 1 µg/ml of specific IgG. High levels of anti-gag antibodies were detected in groups 1, 2, 3, 4, and 7 (Figures 6 and 7). No HIV-1 specific antibodies were detected in groups 5 and 6 (Figure 6).

The ability of the antibodies raised in the immunized monkeys to neutralize HIV-1BX08 virus in human PBMC was assayed based on the reduction of p24 levels.

The neutralization assay was performed essentially as described in reference 18. Briefly, serum dilutions were mixed with HIV-1 BX08 and the mixtures incubated for 1 hour, then added to susceptible human PBMC cells. Titres were recorded as the dilution of serum at which p24 was reduced by 80%. Serum samples were assayed at 1:2, 1:8 and 1:32 dilution on the virus (1:6, 1:24 and 1:26 dilutions after the addition of cells). p24 levels were evaluated by p24-specific ELISA assay.

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[0073] DNA vaccination on its own, group 5, and ALVAC on its own, group 6, had no monkeys showing reduction of p24 levels greater than 80%. The low DNA (600 ug) plus ALVAC, group 4, also showed no monkeys with greater than 80% reduction of p24 titres. VLP plus DNA, either high or low dose (group 1 and 2) showed enhanced reduction of p24 levels compared to VLPs alone, group 7. High dose DNA, group 3, in combination with ALVAC enhanced the ability to elicit p24 or virus neutralising antibodies over the low dose, group 4 or ALVAC alone, group 6. These results indicate that DNA vaccination in combination with VLPs or ALVAC enhanced the levels of virus neutralising antibodies as indicated by the reduction of p24 levels in the sera of the immunized monkeys.

[0074] The percentage reduction of p24 is calculated relative to the amount of p24 produced in the presence of the corresponding dilution of week 2 samples.

SUMMARY OF DISCLOSURE

[0075] In summary of this disclosure, the present invention provides novel immunization procedures and immunogenic compositions for generating virus neutralizing levels of antibodies to a primary HIV isolate and vectors utilized therein and for the generation of components for use therein. Modifications are possible within the scope of this invention.

Table 1 Study Design

Group number	Treatment - Week 0, 4	Treatment – Week 24,44
Rumber	2 ma DV00 DNIA	2 mg DV08 DNIA
	3 mg BX08 DNA	3 mg BX08 DNA
1	50 μg BX08 VLP	50 μg BX08 VLP
	100 μg QS21	100 μg QS21
·	600 μg BX08 DNA	600 μg BX08 DNA
2	50 μg BX08 VLP	50 μg BX08 VLP
·	100 μg QS21	100 μg QS21
3	3 mg BX08 DNA	ALVAC(2) BX08 (1x10 ⁸ pfu)
4	600 μg BX08 DNA	ALVAC(2) BX08 (1x10 ⁸ pfu)
5	3 mg BX08 DNA	3 mg BX08 DNA
6	Control DNA	ALVAC(2) BX08 (1x10 ⁸ pfu)
7	50 μg BX08 VLP	50 μg BX08 VLP
	100 μg QS21	100 μg QS21

Table 2 Number of Monkeys showing > 80% reduction of p24 titre.

Group number	Week 26 Bleed	Week 44 Bleed	
1	3/4	3/4	
2	3/4	4/4	
3	2/4	2/4	
4	0/4	0/4	
5	0/4	0/4	
6 .	0/4	0/4	
7	2/4	3/4	

WO 01/82962 PCT/CA01/00577

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CLAIMS

What we claim is:

1. A method for generating in a host a virus neutralizing level of antibodies to a primary HIV isolate, comprising:

at least one administration of a priming antigen to the host, wherein the priming antigen comprises a DNA molecule encoding an envelope glycoprotein of a primary isolate of HIV-1,

resting the host for at least one specific resting period to provide for clonal expansion of an HIV antigen specific population of precursor B-cells therein to provide a primed host, and

at least one administration of a boosting antigen to the primed host to provide said neutralizing levels of antibodies, wherein the boosting antigen is selected from the group consisting of a non-infectious, non-replicating, immunogenic HIV-like particle having at least the envelope glycoprotein of a primary isolate of HIV-1 and an attenuated viral vector expressing at least an envelope glycoprotein of a primary isolate of HIV-1.

- 2. The method of claim 1 wherein said primary isolate is Bx08.
- 3. The method of claim 2 wherein said DNA molecule is contained in a plasmid vector under the control of a heterologous promoter for expression of the envelope glycoprotein in the host.
- 4. The method of claim 3 wherein the promoter is the cytomegalovirus promoter.
- 5. The method of claim 4 wherein the vector has the identifying characteristics of pCMV3Bx08 shown in Figure 2.
- 6. The method of claim 1 wherein the at least one administration of a priming antigen is at least two administrations of the priming antigen.
- 7. The method of claim 6 wherein the at least one specific resting period is effected after each priming administration.
- 8. The method of claim 1 wherein the at least one specific resting period is between about 2 months to about 12 months.
- 9. The method of claim 1 wherein said non-infectious, non-replicating, immunogenic HIV-like particle comprises an assembly of:
 - (i) an env gene product,

- (ii) a pol gene product, and
- (iii) a gag gene product,

said particle being encoded by a modified HIV genome deficient in long terminal repeats (LTRs) and containing gag, pol and env in their natural genomic arrangement.

- 10. The method of claim 9 wherein the env gene is that from primary isolate BX08.
- 11 The method of claim 1 wherein said non-infectious, non-replicating, immunogenic HIV-like particle is administered in conjunction with an adjuvant.
- 12. The method of claim 11 wherein the adjuvant is QS21.
- 13. The method of claim 1 wherein said attenuated viral vector is an attenuated avipoxvirus
- 14. The method of claim 13 wherein the attenuated viral vector contains a modified HIV-genome deficient in long terminal repeats, wherein at least the *env* gene is that from primary isolate BX08.
- 15. The method of claim 14 wherein the attenuated avipoxvirus vector is the attenuated canary poxvirus ALVAC.
- 16. The method of claim 15 wherein the attenuated canary poxvirus vector has the identifying characteristics of vCP1579.
- 17. The method of claim 1 wherein the at least one administration of a boosting antigen is at least two administrations of a boosting antigen.
- 18. A vector, comprising a DNA sequence encoding an envelope glycoprotein of a primary isolate of HIV-1 under the control of a heterologous promoter for expression of the envelope glycoprotein in a host organism.
- 19. The vector of claim 18 wherein the vector is a plasmid vector.
- 20. The vector of claim 18 wherein said primary HIV-1 isolate is Bx08.
- 21. The vector of claim 20 wherein the promoter is the cytomegalovirus promoter.
- 22. The vector of claim 21 which has the identifying characteristics of pCMV3Bx08 shown in Figure 2.
- 23. The vector of claim 18 wherein the vector is an attenuated viral vector.

- 24. The vector of claim 23 wherein the attenuated viral vector is a attenuated avipoxvirus vector.
- 25. The vector of claim 24 wherein the attenuated avipoxvirus vector is the attenuated canary poxvirus vector ALVAC.
- 26. The vector of claim 25 wherein the attenuated viral vector has the identifying characteristics of vCP1579 shown in Figure 4.
- 27. A vector, comprising a modified HIV genome deficient in long terminal repeats and a heterologous promoter operatively connected to said genome for expression of said HIV genome in mammalian cells to produce non-infectious, non-replicating and immunogenic HIV-like particles, wherein at least the *env* gene is that from a primary isolate of HIV-1.
- 28. The vector of claim 27 wherein the vector is a plasmid vector.
- 29. The vector of claim 28 wherein the primary HIV-1 isolate is BX08.
- 30. The vector of claim 29 wherein the promoter is type IIA metallothionein promoter.
- 31. The vector of claim 30 which has the identifying characteristics of p133B1 shown in Figure 3.

Figure 1 Plasmid pCMV.Bx08.gp160

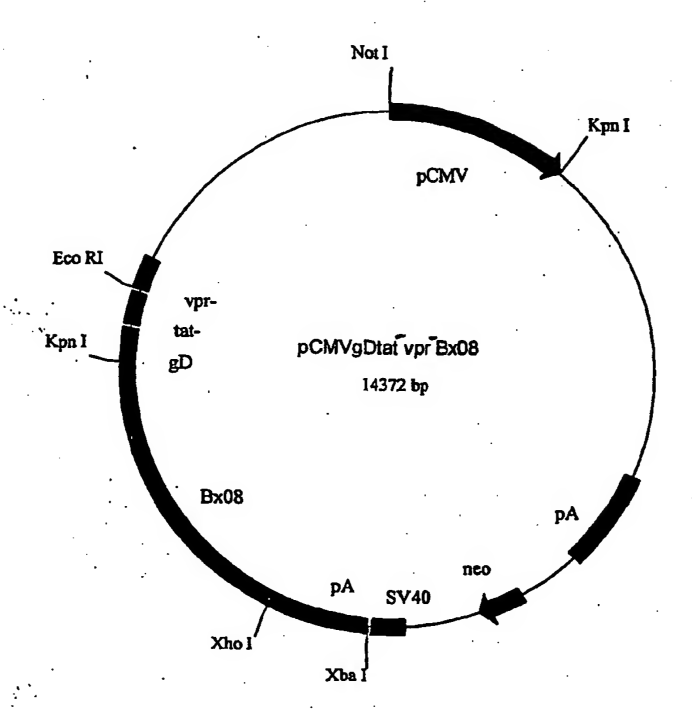


Figure 2 DNA immunization plasmid pCMV3Bx08.

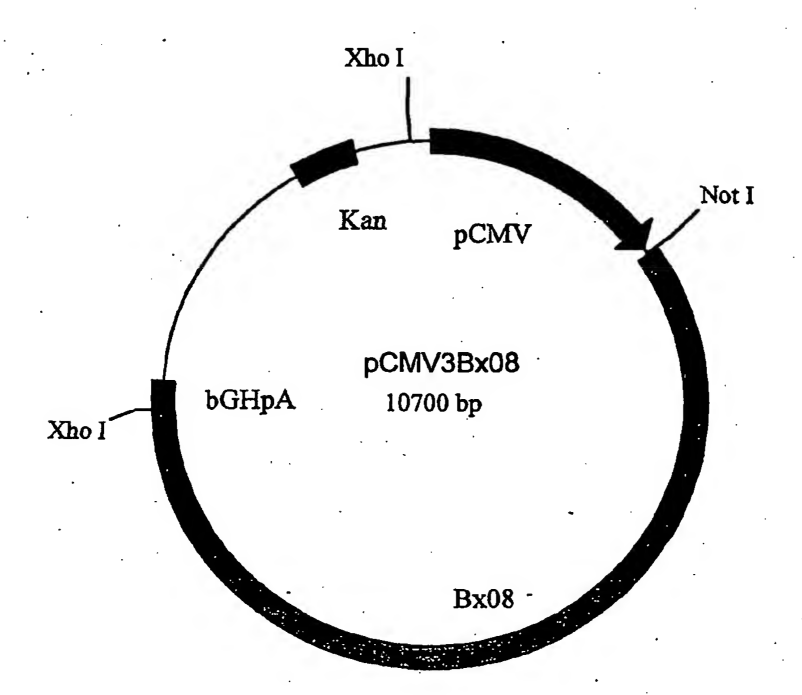


Figure 3. Pseudovirion Expression Plasmid p133B1 HIV-1 Bx08

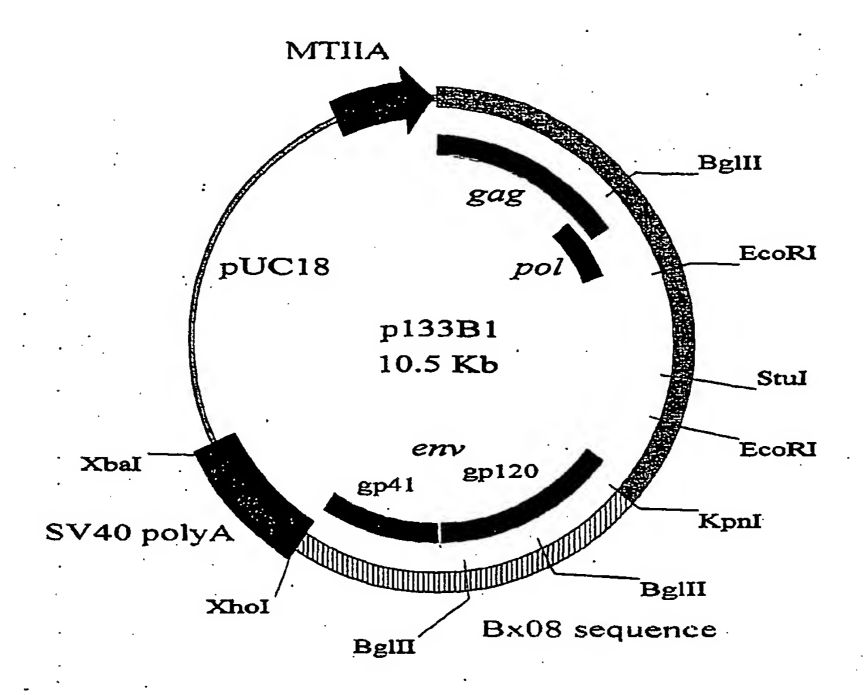
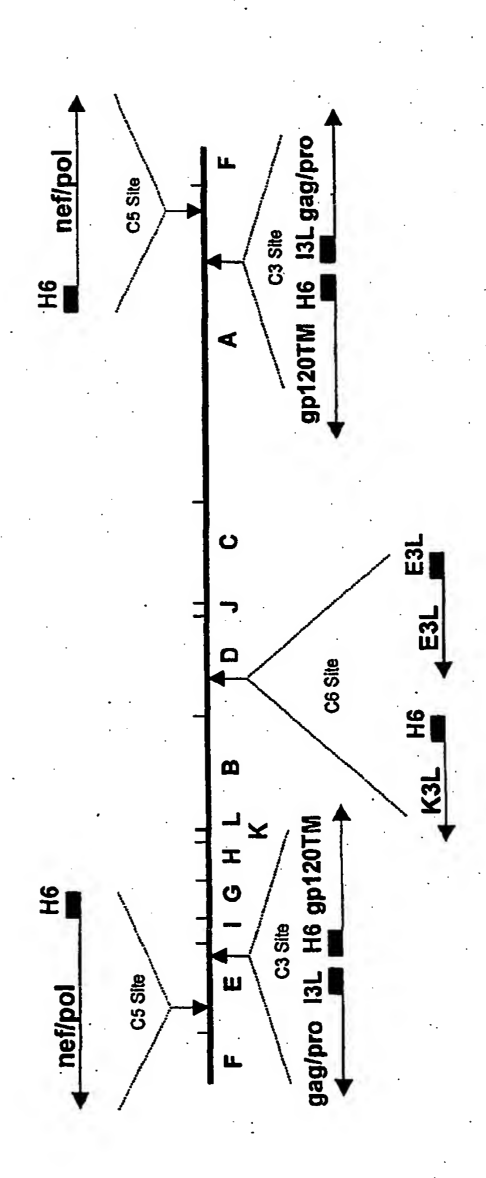
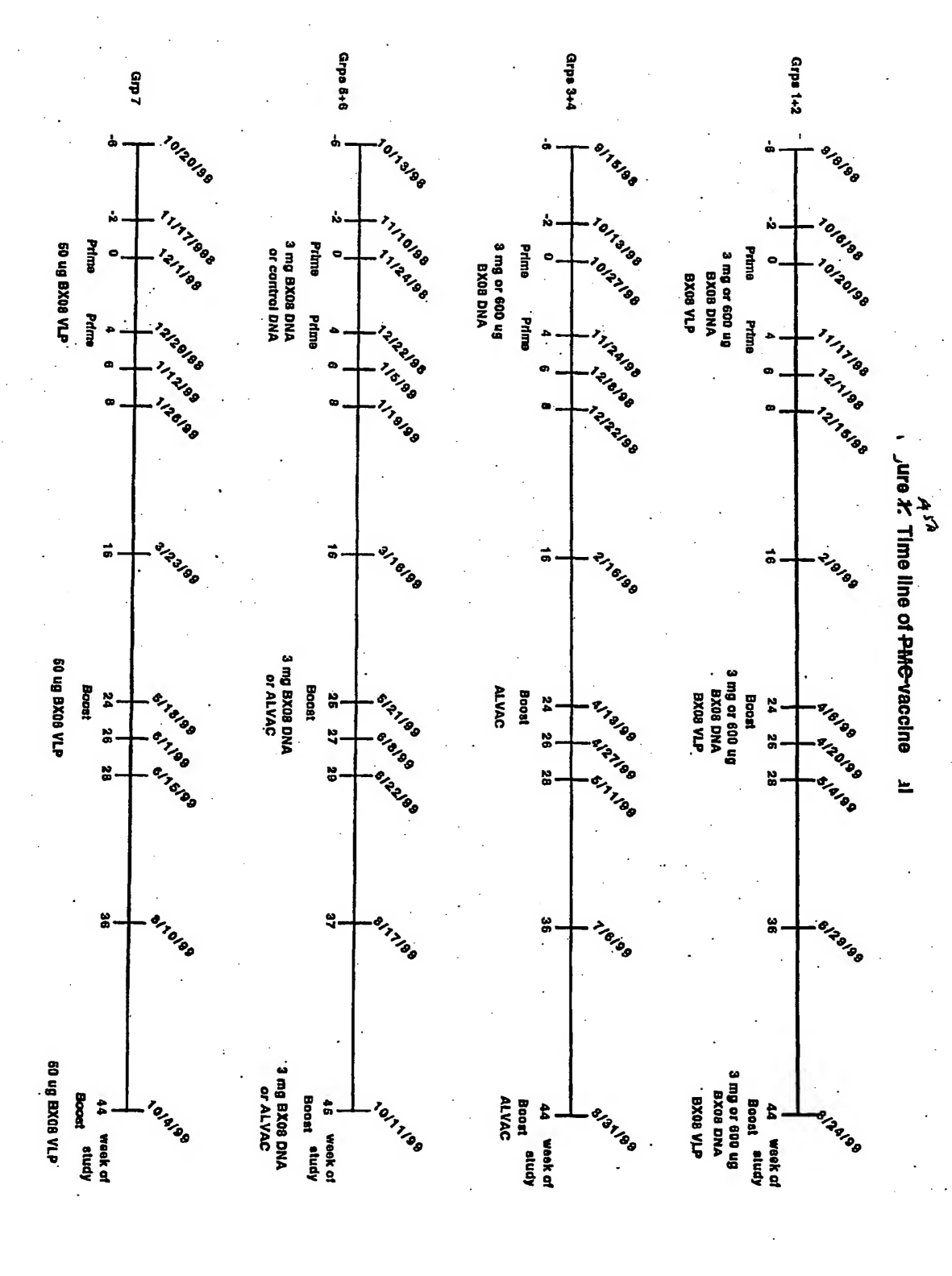


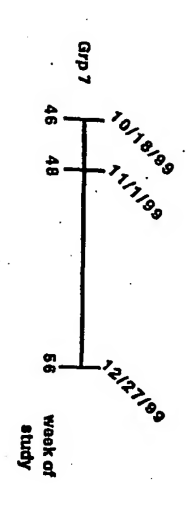
FIGURE 4

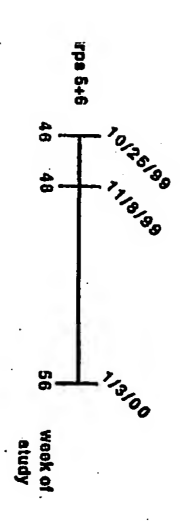
ALVAC(2)120(BX08)GNP (vCP1579)

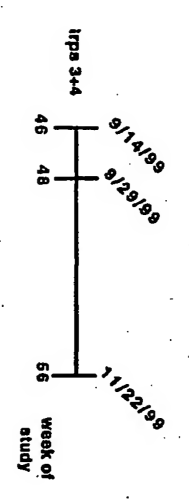
ALVAC Xhol Restriction Map)

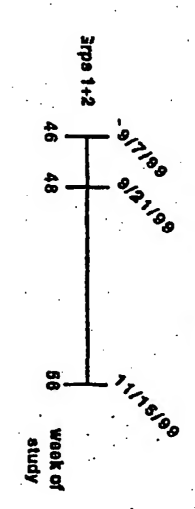


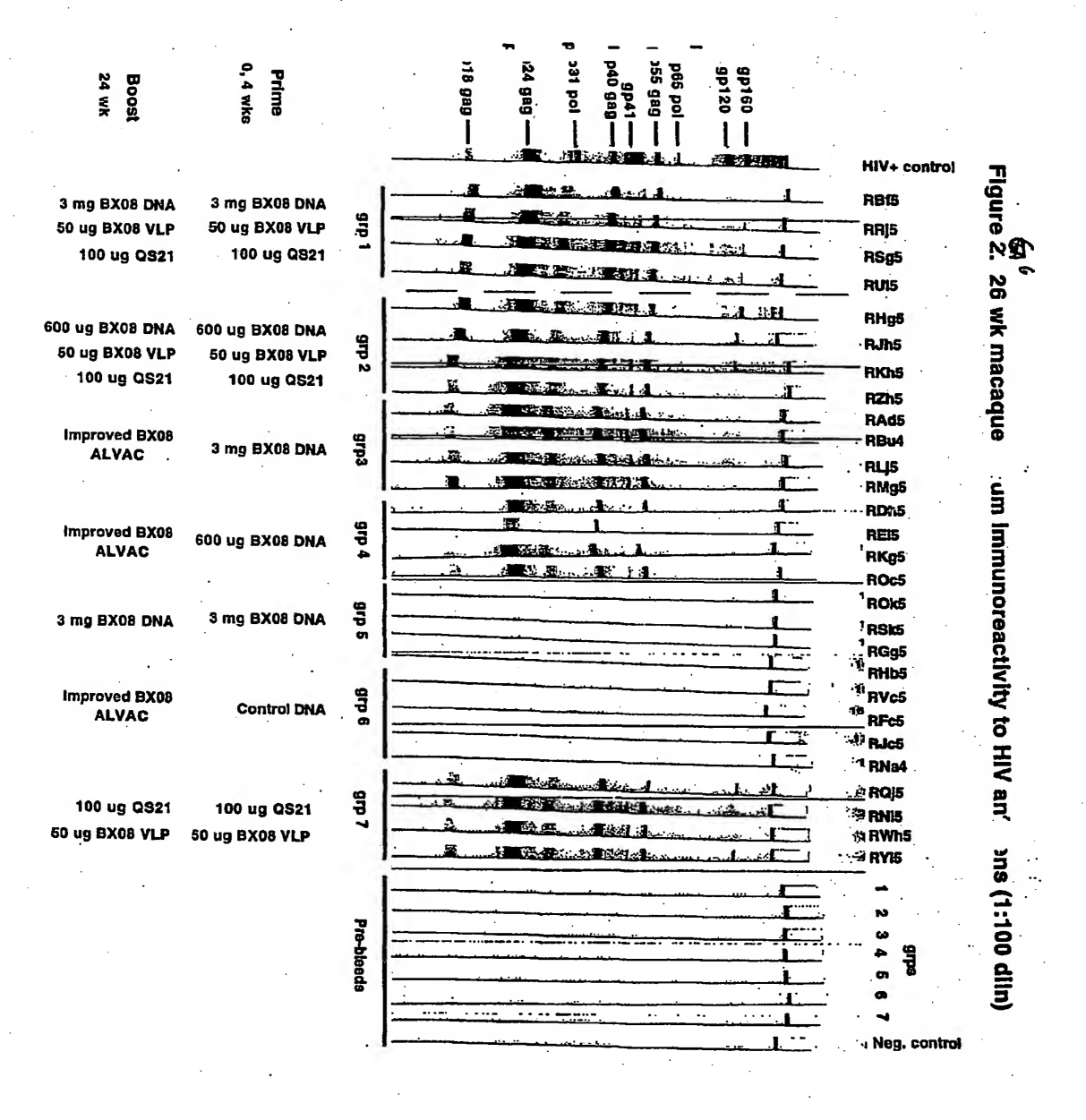




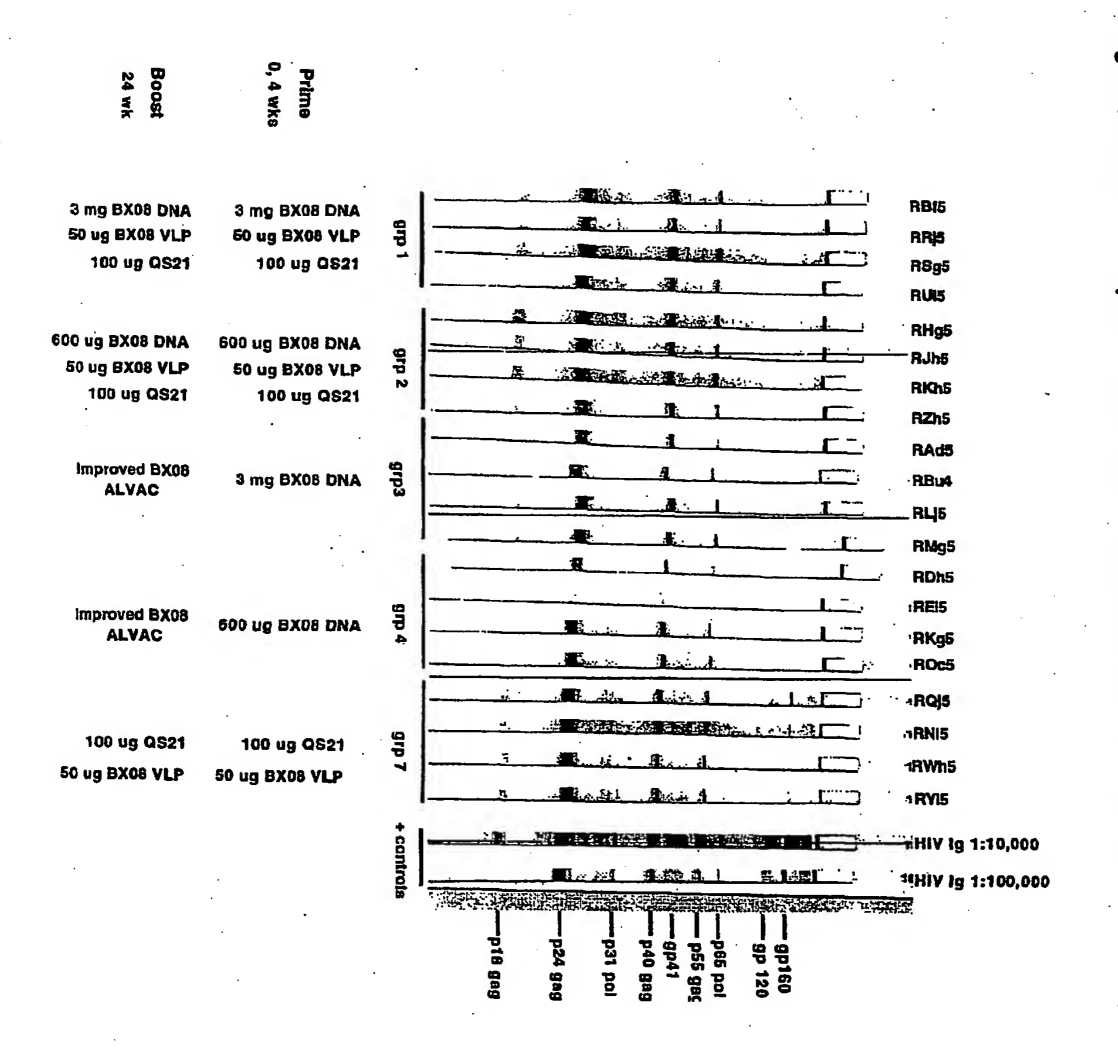


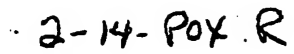


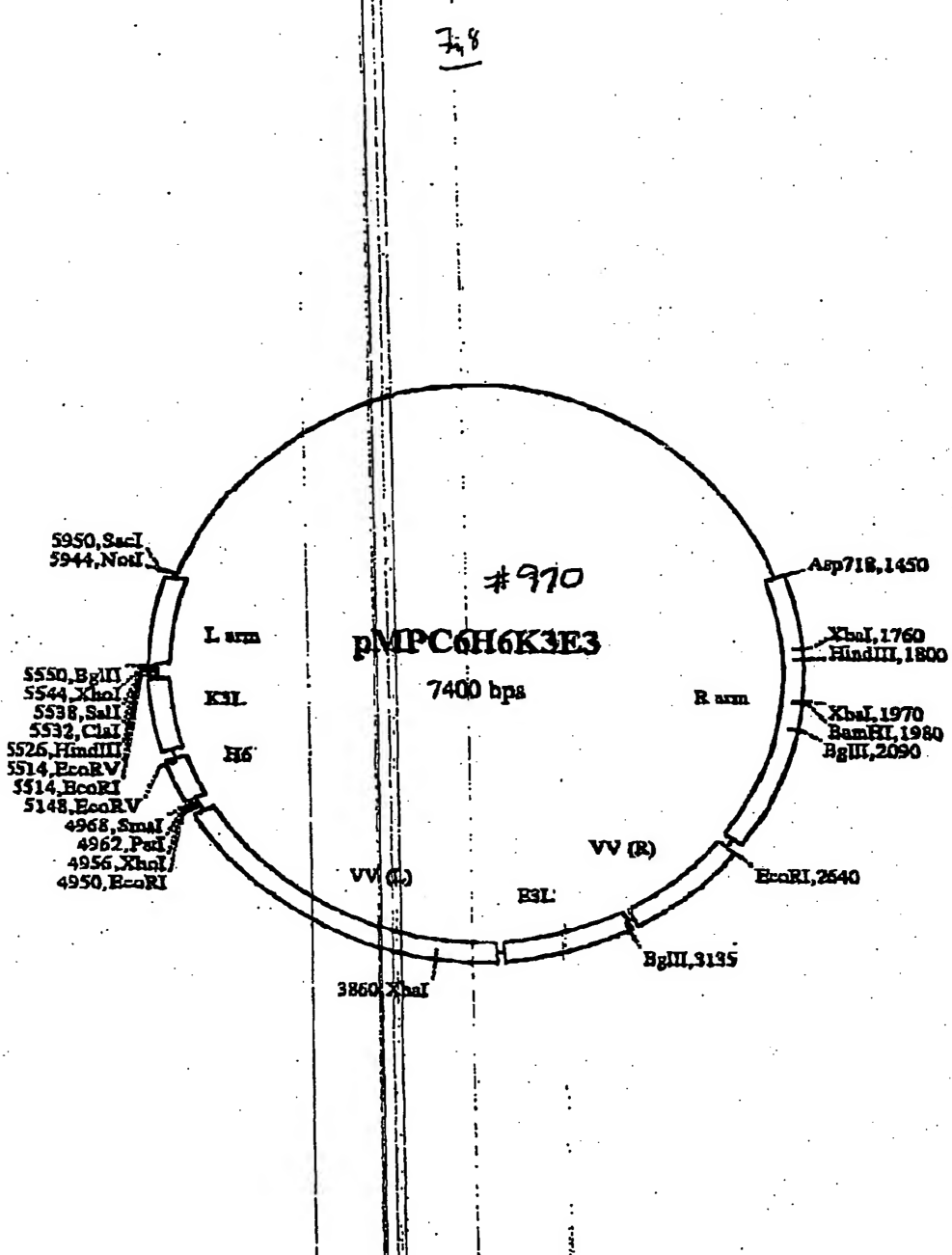












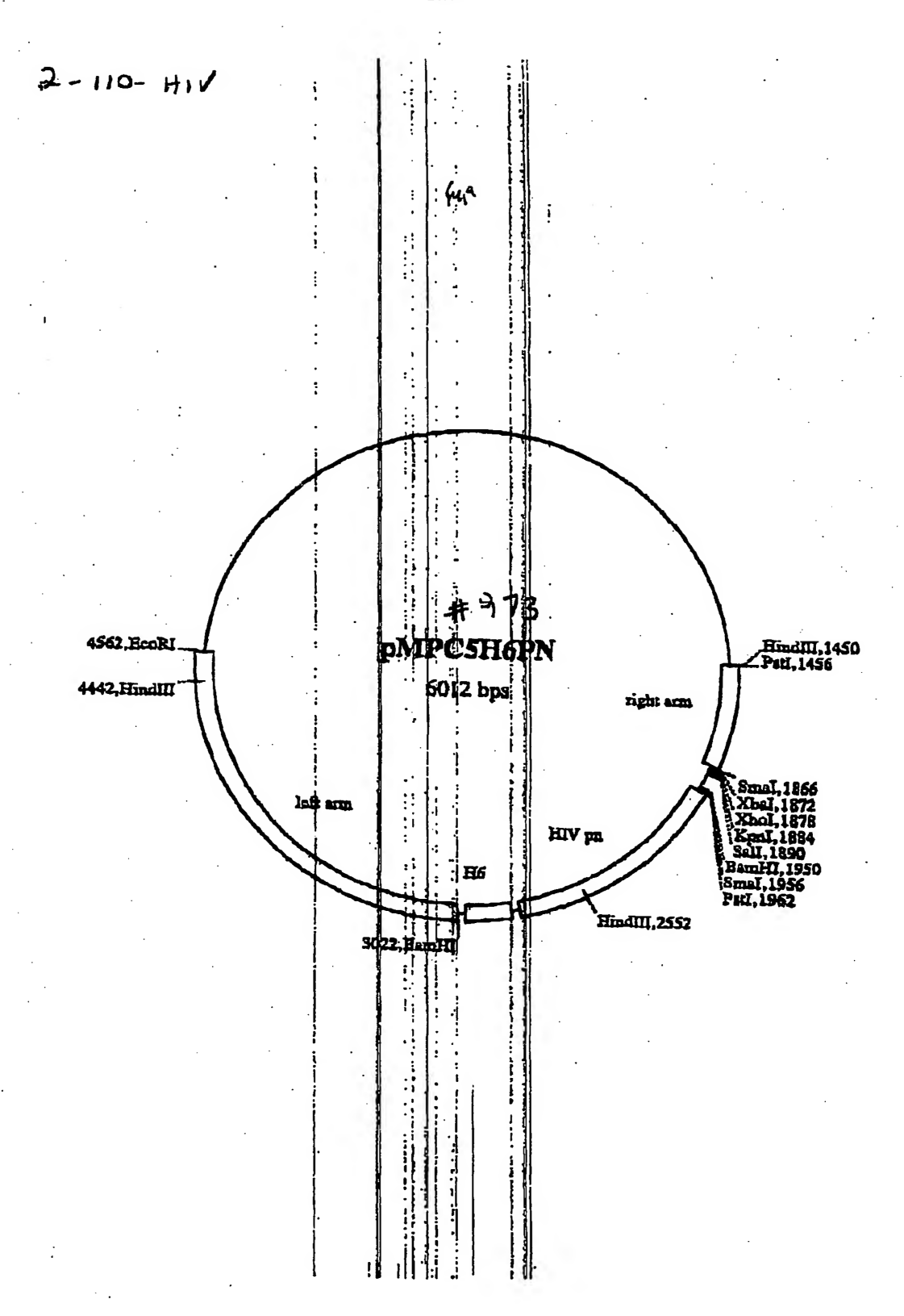


Figure 10 Plasmid pHIV76

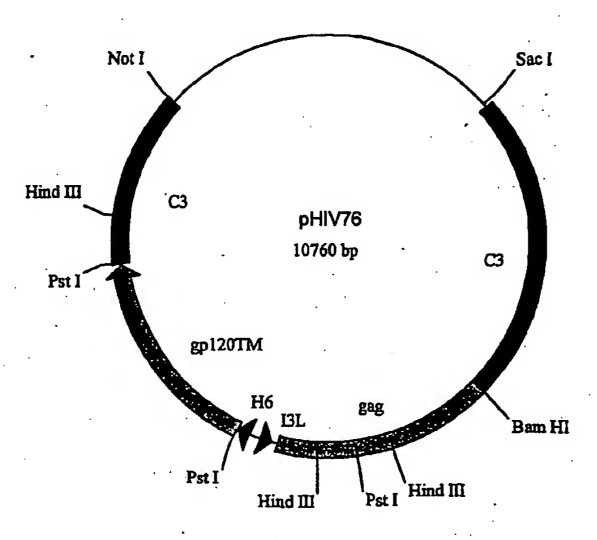


Figure 11 vCP1579: H6/HIV Pol/Nef epitope cassette in ALVAC C5 site

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1 TTTTTTTCAT TATTTAGAAA TTATGCATTT TAGATCTTTA TAAGCGGCCG TGATTAACTA
  61 GTCATAAAAA CCCGGGATCG ATTCTAGACT CGAGGGTACC GGATCTTAAT TAATTAGTCA
 121 TCAGGCAGGG CGAGAACGAG ACTATCTGCT CGTTAATTAA TTAGGTCGAC GGATCCCCCA
 181 ACAAAAACTA ATCAGCTATC GGGGTTAATT AATTAGTTAT TAGACAAGGT GAAAACGAAA
 241 CTATTTGTAG CTTAATTAAT TAGAGCTTCT TTATTCTATA CTTAAAAAGT GAAAATAAAT
301 ACAAAGGTTC TTGAGGGTTG TGTTAAATTG AAAGCGAGAA ATAATCATAA ATTATTTCAT
 361 TATCGCGATA TCCGTTAAGT TTGTATCGTA ATGCCACTAA CAGAAGAAGC AGAGCTAGAA
 421 CTGGCAGAAA ACAGAGAGAT TCTAAAAGAA CCAGTACATG GAGTGTATTA TGACCCATCA
481 AAAGACTTAA TAGCAGAAAT ACAGAAGCAG GGGCAAGGCC AATGGACATA TCAAATTTAT
 541 CAAGAGCCAT TTAAAAATCT GAAAACAGGA ATGGAGTGGA GATTTGATTC TAGATTAGCA
601 TTTCATCACG TAGCTAGAGA ATTACATCCT GAATATTTTA AAAATTGTAT GGCAATATTC
661 CAAAGTAGCA TGACAAAAAT CTTAGAGCCT TTTAGAAAAC AAAATCCAGA CATAGTTATC
 721 TATCAATACA TGGATGATTT GTATGTAGGA TCTGACTTAG AAATAGGGCA GCATAGAACA
781 AAAATAGAGG AGCTGAGACA ACATCTGTTG AGGTGGGGAC TTACAACCAT GGTAGGTTTT
841 CCAGTAACAC CTCAAGTACC TTTAAGACCA ATGACTTACA AAGCAGCTGT AGATCTTTCT
901 CACTTTTTAA AAGAAAAAGG AGGTTTAGAA GGGCTAATTC ATTCTCAACG AAGACAAGAT
961 ATTCTTGATT TGTGGATTTA TCATACACAA GGATATTTTC CTGATTGGCA GAATTACACA
1021 CCAGGACCAG GAGTCAGATA CCCATTAACC TTTGGTTGGT GCTACAAGCT AGTACCAATG
1081 ATTGAGACTG TACCAGTAAA ATTAAAGCCA GGAATGGATG GCCCAAAAGT TAAACAATGG
1141 CCATTGACAG AAGAAAAAT AAAAGCATTA GTAGAAATTT GTACAGAGAT GGAAAAGGAA
1201 GGGAAAATTT CAAAAATTGG GCCTTAATTT TTCTGCAGCC CGGGGGATCC TTTTTATAGC
1261 TAATTAGTCA CGTACCTTTG AGAGTACCAC TTCAGCTACC TCTTTTGTGT CTCAGAGTAA
1321 CTTTCTTTAA TCAATTCCAA AACAG
```

Upstream (right) flanking sequence: 1-266

VV H6 promoter: 267-390

HIV pol/nef/pol/nef/pol cassette: 391-1227

Downstream (left) flanking sequence: 1227-1345

Figure 12
E3L and K3L genes in C6

	10	20		40	50	60	70	80	90	100	110
•	GAGCTOGCGG CTCGAGCGCC	CCGCCTATCA GGCGGATAGT	AAAGTCTTAA	TGAGTTAGGT ACTCAATCCA	GTAGATAGTA	TAGATATTAC	TACAAAGGTA	TTCATATTTC	CTATCAATIC CATAGTTAAG	TAAAGTAGAT (ATTTCATCTA (ATAATTATA TATTAATAT
	120	130	140	. 150	160	170	180	190	200	210	220
	ACTCAAAGAT TGAGTTTCTA	GATGATAGTA CTACTATCAT	GATAATAGAT CIATIATCIA	ACGCTCATAT TGCGAGTATA	AATGACTGCA TTACTGACGT	AATTTGGACG TTAAACCTGC	GTTCACATTT CAAGTGTAAA	TAATCATCAC ATTAGTAGTG	GCGTTCATAA CGCAAGTATT	GITTCAACTG (CAAAGITGAC (CATAGATCAA GTATCTAGTT
	230	240			270	280	290	300	310	320	330
	AATCTCACTA TTAGAGTGAT	AAAAGATAGC TTTTCTATCG	CGATGTATTT CCTACATAAA	GAGAGAGATT CTCTCTCTAA	GGACATCIAA CCTGTAGATT	CTACGCTAAA GATGCGATTT	GAAATTACAG CITTAATGTC	ATATAAATAA TATTTATTAA	TACATAATGG ATGTATTACE	ATTTTGTTAT TAAAACAATA	CATCAGITAT STAGTCAATA
	340	350		• •	380		400	410	• •	430	440
	ATTTAACATA TAAATTGTAT	AGTACAATAA TCATGTTATT	AAAGTATTAA TITCATAATT	ATAAAATAC TATTTTTATG	TTACTTACGA AATGAATGCT	AAAAATGACT TTTTTACTGA	AATTAGCTAT TTAATCGATA	AAAAACCCAG	ATCTCTCGAG TAGAGAGCTC	GTCGACGGTA CAGCTGCCAT	TCGATAAGCT AGCTATTCGA
	450	460	•	470	480	490	500	•	10	520	530
	TGATATCGAA ACTATAGCTT	TTCATAAAA AAGTATTTT	AA T AAC T	AC AGA TOT	TTA GGA AAA M· R K	CAT TAX CITY Y N V	TAG ATA TA	AT AGG AAA I	ACA TAT TAG T Y D	AAC TOT AAT TIG AGA TIA V R I	GTG AAA V K
	540	. 5:	50	560	570	580	590		600.	610	620
	AAC TIT TAC TIG AAA ATC <v k="" td="" v<=""><td>TCA AAA G</td><td>GG ATG GTC</td><td>TTT ATC CCT AAA TAG GGA K D R</td><td>TAT ANG TTO</td><td>TAT AGA T</td><td>C CAT ATG</td><td>CAT CTT AAC</td><td>ACT CTC TO</td><td>CAA GAT AG</td><td>C TTC AGA</td></v>	TCA AAA G	GG ATG GTC	TTT ATC CCT AAA TAG GGA K D R	TAT ANG TTO	TAT AGA T	C CAT ATG	CAT CTT AAC	ACT CTC TO	CAA GAT AG	C TTC AGA
							68		690	700	710
	CTG AGG ATI	GTC AAA A	40 • • • AG ATA AAT (650 GTA TAG AGC	ATA ATC CTT	670 T CTC GTA T	LC TCT GCC (CTT TAT TAC	ATC GCC CG	C ATT GGG CA	A CGA ATA
	CAC TCC TAT	CAG TIT T	TC TAT TTA	CAT ATC TCG	TAT TAG GAN	GAG CAT A	TG AGA COOG (EAA ATA ATG	TAG CGG GC	N P I	T GCT TAT
	720	7:	30 7	40 7!	50 76		70 7	-		00 83	.0
		•	• •					• •	• •	• •	•
	TGT TTT ACC	TTC GTA TO	CG ATACAAAC	TT AACOGATA	TC GCGATAATO	SA AATAATIT	AT GATTATTT	CT CGCTTTCA	AT TTAACACA	AC CCTCAAGAI TG GGAGTTCT	ug 70
	TOT TIT ACC	ARG CRT AG TTC GTA TO L M ·	CG ATACAAAC GC TATGITTG	TT AACOGATA: AA TTGCCTAT	TC GCGATAATC	SA AATAATIT	AT GATTATTY EA CTAATAAA	CT CGCTTTCA GA GCGAAAGT 890	AT TTAACACA TA AATTGTGT	AC CCTCAAGAI TG GGAGTTCT	
	TGT TTT ACC <c 820="" a="" ctttgtattt<="" f="" td=""><td>AAG CAT AG TTC GTA TO L M - K3L B30</td><td>EG ATACANAC GC TATGITIG 840 TITNAGTATA</td><td>TT AACOGATA: AA TTGCCTATE 850 GAATAAGAA</td><td>GCGATAATC AG CGCTATTAC 860 AGCTCTAATT</td><td>SA AATAATITE TTATTAAN E70 AATTAATGAA</td><td>AT GATTATTO</td><td>CT CGCTTTCA GA GCGAAAGT 890 CGTTTTCCCC</td><td>AT TTAACACA TA AATTGTGT 900 TTGGCGTATC</td><td>AC CCTCAAGAI TG GGAGTTCT 910 ACTAATTAAT</td><td></td></c>	AAG CAT AG TTC GTA TO L M - K3L B30	EG ATACANAC GC TATGITIG 840 TITNAGTATA	TT AACOGATA: AA TTGCCTATE 850 GAATAAGAA	GCGATAATC AG CGCTATTAC 860 AGCTCTAATT	SA AATAATITE TTATTAAN E70 AATTAATGAA	AT GATTATTO	CT CGCTTTCA GA GCGAAAGT 890 CGTTTTCCCC	AT TTAACACA TA AATTGTGT 900 TTGGCGTATC	AC CCTCAAGAI TG GGAGTTCT 910 ACTAATTAAT	
	TGT TTT ACC <c 820="" a="" ctttgtattt<="" f="" td=""><td>ANG CRT AND THE STORY OF THE ST</td><td>840 TITLAGTATA AAATTCATAT</td><td>AA TIGOCTATA 850 GAATAAAGAA CITATICIT 960</td><td>GCGATAATO AG CGCTATTAO 860 AGCTCTAATT TCGAGATTAA</td><td>SA AATAATITE TTATTAAN E70 AATTAATGAA</td><td>AT GATTATTO FA CTARTAAA 880 CAGATTGTTT GTCTAACAAA</td><td>CT CGCTTTCA GA GCGAAAGT 890 CGTTTTCCCC GCAAAAGGGG</td><td>900 TIGGCGTATC</td><td>910 ACTANTANT TGATTANTA</td><td>920 TAACCOGGC ATTGGGCCCG</td></c>	ANG CRT AND THE STORY OF THE ST	840 TITLAGTATA AAATTCATAT	AA TIGOCTATA 850 GAATAAAGAA CITATICIT 960	GCGATAATO AG CGCTATTAO 860 AGCTCTAATT TCGAGATTAA	SA AATAATITE TTATTAAN E70 AATTAATGAA	AT GATTATTO FA CTARTAAA 880 CAGATTGTTT GTCTAACAAA	CT CGCTTTCA GA GCGAAAGT 890 CGTTTTCCCC GCAAAAGGGG	900 TIGGCGTATC	910 ACTANTANT TGATTANTA	920 TAACCOGGC ATTGGGCCCG
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	2780	• 1	790	*	•	283	•	282	•	2830		2840		2850		2850		
																		GTCATCTATA
		GILLIGTTA	III T	CHILITEC	AT G	MAATAGT	AA AA	ARATANG.	r AGT	MGAGAC	CACCA	AUCAG	CAAAG	MIAGC	TTAC	NI CUAG	ALTARTTGG	CAGTAGATAT
	GATTOTOTIG	U, U, I															•	
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	2890		900	29		292	£0	2931	•	2940		2950		2960	•	2970	2980	2990
	2890	. 2	٠	•	•	•	•	• ,	• .	•	•	•	•	•		• •	•	•
	2890 GGIGATGCTG	2 • GITCTGG	• •GA T	* TCT9GAG	GA G	• ATGGATT	• IT TA	TETGGAN	TAA S	CICIGII	ATTIC	c rici	TTTCA	* TATOI	CGAT	rgcgit	GTAÄCATTAI	CATTGCGALA
	2890 GGIGATGCTG	2 • GITCTGG	• •GA T	* TCT9GAG	GA G	• ATGGATT	• IT TA	TETGGAN	TAA S	CICIGII	ATTIC	c rici	TTTCA	* TATOI	CGAT	rgcgit	GTAÄCATTAI	•
	2890 GGIGATGCTG (CCACTACGAC (2 TTCTGG CANGACC	AGA T	* TCTGGAG AGACCTG	GA G	ATGGATT/ TACCTAAT	TA AT	TCTGGAA AGACCTT	Z AAT Z TIA	CTCTGTT GAGACAA	ATTIC	CTTIGT GAACA	TITCA'	IGTAT ACATA	CGAT	rgcgit ACGCAA	GTAACATTAI CATTGTAATT	CATTGCGÀA CTAACGCTIT
	2890 GGIGATGCTG	2 TTCTGG CANGACC	• •GA T	* TCTGGAG AGACCTG	* GA G CT C	• ATGGATT	TA AT	TETGGAN	Z AAT Z TIA	CICIGII	ATTIC	CTIGI GAACA 3060	TITCA ARAGT	* TATOI	CGAT	rgcgit	GTAACATTAI CATTGTAATT	CATTGCGAAA CTAACGCTIT
	2890 GGIGATGCTG (CCACTACGAC (3000 TGCTCTAAAT 7	2 TTCTGG CAAGACC	AGA TICT A	TCTGGAG IAGACCTC 30	GA G CT C	ATGGATTI TACCTAAT 303	TA AT	TCTGGAAAAAGACCTTY	AAT TIA	CTCTGTT GAGACAA 3050 TAACTAG	ATTTC TAXAG	CTTOT GAACA 3060 GTTCG	TTTCA' ARAGTI	TGTATACATA	CGAT	IGCGIT ACGCAA 3080	GTAACATTAI CATTGTAATT 3090	CATTGCGÁA CTAACGCTIT 3100 CTAGTATICC
	2890 GGIGATGCTG (CCACTACGAC (3000 TGCTCTAAAT 7	2 TTCTGG CAAGACC	AGA TICT A	TCTGGAG IAGACCTC 30	GA G CT C	ATGGATTI TACCTAAT 303	TA AT	TCTGGAAAAAGACCTTY	AAT TIA	CTCTGTT GAGACAA 3050 TAACTAG	ATTTC TAXAG	CTTOT GAACA 3060 GTTCG	TTTCA' ARAGTI	TGTATACATA	CGAT	IGCGIT ACGCAA 3080	GTAACATTAI CATTGTAATT 3090	CATTGCGAAA CTAACGCTIT

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	3110	3120	3130	3140	3150	3160	3170	3180	3190	3200	3210
										TGTATTCTAC ACATAAGATG	
	ra designation	CALCETOIO	CONCATANCE	·	·		resilent g	·	MIMMINIA	ACKIANOMIO	MULTURALIA.
	3220	3230	3240	3250	3260	3270	3280	3290	3300	3310	3320
										TAAATCATAT ATTTAGTATA	AATAATGAAA
	INCOINMAI	AISOCIIAII	CICIAICUCI	IÇCIIAMAN	·	1101104011	·	MILLOCITI	TICKTITIAL	AIIIAGIAIA	IIMIIMCIII
	3330	3340	3350	3360	3370	3380	3390	3400	3410	3420	3430
	• •	• •		• •		• •	•		• •	GCAAATACAG	* *
										CGTTTATGTC	
	3440	3450	3460	3470	3480	3490	3500	3510	3520	3530	3540
	•	• •		• •	• •		• •	• •	• •	ATAATATAAT	
											TTTCTAAGTA
	3550	3560	3570	3580	3590	3600	3610	3620	3630	3640	3650
	• •	• •	• •	• •	• •	• •	• •	• •		ATTAAAATAG	• •
										TAATTTTATC	
	3660	3670	3680	3690	3700	3710	3720	3730	3740	3750	3760
	AGATGTAAAT	AATTATTTGG	AGGTAAAGGA	• •	• •	CACATGGAAA		• •			GGATTTACAG
										CTTAAAAAAT	
	3770	3780	3790	3600	3810	3820	3830	3840	3850	3860	3870
										TAATAGCCAG	
	CACAATATAC	atagpigtta	TGTCCGTCTA	CATACCAATA	CCATTTTGTG	ACATTGCCCT	TCGTCGTAAG	ATACCATTGA	CCGGATACAA	ATTATEGGTC	TACTAAAATC
	3880	3890	3900	3910	392 ò	3930	3940	3950	3960	3970	3980
											CTACTAATAA
	AGATATTTGT	AAAATGGTGT	TTATTATCCT	AGGAGATCTA	AATATTAAAT	TATAGATIGT	TGITGITTI	TTAAATTGCT	ACATACCGGT	CTTCATAAAA	GATGATTATT
	3990	4000		4020	4030				4070		
				–							TITATATIAA
	TCIATITCIA	TCAGATAGAA	TAGATGTTCT	ATACTTTCTT	CTATTAGTAA	ATCATCATCG	ATGATTATAC	CITICITIAC	ATATGITTIT	GCACCTTCGA	AAATATAATT
	4100	4110	4120	4130	4140	4150	4160		4180	4190	
											CAGTGATATA
	TATEGTATAA	TGATCTTCTA	ARTITIAGAT	CIGAATCATA	TIGITITETC	AATTTACGGT	TATAGCTAAG	ATATAAAGTA	GIATIGICAT	CATGTAATTA	GTCACTATAT
	4210	4220	4230	4240	4250	4260	4270	4280	4290	. 4300	4310
		CTACAGACTC	AACTATGCAA		ATATGCCAAT			TAGAACTAAA			ATAGGATACG
	GACITTUCIA	CATOTOTORS	TTGATACGIT	CETTATTCGT	TATACGGITA	ATACAGATTA	TAAAATTGAA	ATCITGATIT	TGCAAGATGG	TTATGATTTT	TATCCTATGC
	4320	4330	4340	4350	4360	4370	4380	4390	. 4400	4410	4420
		TTANAAGCTG			GAAGAAATAC						CAAACTIGIA
	HC LATTUCEAC	DALLITIAN	STATTATE	ATTUCTACAT	CTICTTIATG	ARACAAGATA	TOGRADICETC	CTTTCTTGAA	ATCTIGITGA	ATTCAAATTA	GTTTGAACAT

4430

TITATGAAGG TACC AAATACTTCC ATGG

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- with international search report
- (88) Date of publication of the international search report: 21 March 2002

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



(54) Title: IMMUNIZING AGAINST HIV INFECTION

(57) Abstract: A virus neutralizing level of antibodies to a primary HIV isolate is generated in a host by a prime-boost administration of antigents. The primary antigen is a DNA molecule encoding an envelop glycoprotein of a primary isolate of HIV-1 while the boosting antigen is either a non-infectious, non-replicating HIV-like particle having the envelope glycoprotein of a primary isolate of HIV-1 or an attenuated viral vector expressing an envelope glycoprotein of a primary isolate of HIV-1.

INTERNATIONAL SEARCH REPORT

PC1/CA 01/00577

A. CLASSIFICATION OF SUBJECT MATTER-IPC 7 A61K39/21 A61K39/39 C12N15/86 CO7K14/16 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) A61K C07K C12N IPC 7 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, PAJ, BIOSIS C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. 1,8,18, CAVER T E ET AL: "A novel vaccine regimen utilizing DNA, vaccinia virus and protein 19 immunizations for HIV-1 envelope presentation" VACCINE, BUTTERWORTH SCIENTIFIC. GUILDFORD, GB, vol. 17, no. 11-12, March 1999 (1999-03), pages 1567-1572, XP004158286 ISSN: 0264-410X the whole document 2-7 9-17Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents : *T* later document published after the international filing date or priority date and not in conflict with the application but *A* document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance 'E' earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannol be considered to involve an inventive step when the *O* document referring to an oral disclosure, use, exhibition or document is combined with one or more other such documents, such combination being obvious to a person skilled other means in the art. document published prior to the international filing date but *&* document member of the same patent family later than the priority date claimed Date of mailing of the international search report Date of the actual completion of the international search 02/01/2002 13 December 2001 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Galli, I Fax: (+31-70) 340-3016

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INTERNATIONAL SEARCH REPORT

Inter 'ional Application No PCI/CA 01/00577

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